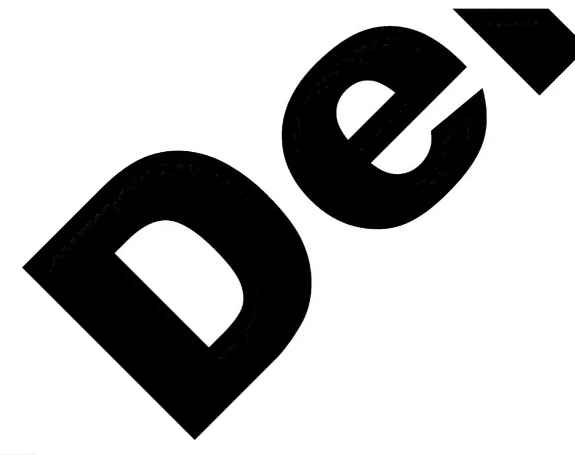
# Approved For Release STAT 2009/08/26 :

CIA-RDP88-00904R000100110



Approved For Release 2009/08/26 :

CIA-RDP88-00904R000100110



Third United Nations
International Conference
on the Peaceful Uses
of Atomic Energy

A/CONF.28/P/766 UKRAINIAN SSR May 1964 Original: RUSSIAN

Confidential until official release during Conference

DETERMINATION OF INTENSITY OF RADIOACTIVE CONTAMINATION

IN THE OCEAN BASED ON THE NEW DATA OF EXCHANGE PROCESS.

V.I.Belyaev, A.G.Kolesnikov, B.A. Nelepo.

Marine Hydrophysical Institute of the Academy of Sciences of the UkssR.Sevastopol.U S S R.

25 YEAR RE-REVIEW

### 71. INTRODUCTION.

Numerous works encountered in scientific literature of recent years show that some authors are very much interested in the problem of evaluation of spreading rate of radioactive contamination in seas and oceans. It is known that this question requires profound knowledge of intensity of processes of the water mass mixing. Inverse problems are also known, that is application of data of radioisotope distribution with depth for determination of intensity of mixing. In them conclusion of intensity of process of mixing made is based on the analysis of the vertical distribution of radioisotopes deposited on the surface layer of the ocean after nuclear weapon tests (as a result of fall- out, in general, of stratospheric character). Some investigators use for this purpose data of distribution of the natural radioactive elements of coangomical origin, such as T.C-14.Be-7.

In the first case the term " age " was introduced to evaluate intensity process of spreading of radioactive contamination. Later this term was broadly used by various investigators for solving some practical problems. Those investigators sho studied radiocarbon distribution in ocean waters determined the age of the Atlantic Ocean to be about 500 years. For the radioactive products in the upper mixed layer was determined as 30-100 years.

Bowen and Sugihara / 1 / are believed to be the first to receive experimental data, which disproved such meaning of the intensity of deep water contamination in the ocean. In \$357 they discovered penetration of radiomelides to the depth of 1000 m. The age of these radiomelides could not be greater than three years, because the intensive fall-out of radioactive isotopes began after 1954. Further observations carried out by Miyake and others /2/confirmed the considerable penetration of radioactive cs and Sr to much greater depths in the ocean. Thus the results of analysis of radioactive isotope distribution in the ocean show that values obtained by radiocarbon technique are wrong. In fact, the term "age" (though successful in geology) could not be mechanically transfered to ocean waters and its usage supposing that the ocean waters did not mix and were stable — was wrong.

Rocco and Broscher noting many contradictions which could be encountered in modern knowledges of processes of the vertical radioactivity transfer tried to make a control experiment. But they received little material and it differed from the well corroborated knowledges of concentrations of radioactive Sr and Cs at intermediate and great depths in the ocean. As so, it is necessary to make some additional experiments and to analyze the obtained data with great care.

on the other hand it exists now the intensity of vertical mixing using data of radioisotope distribution which is in many cases based on Fikkian equation of diffusion.

Some works are known now in which the suthers made an attempt to calculate by indirect method values of turbulent diffusivity in the ocean evaluating intensity of the vertical transfer some simplifying suppositions were made. In particular, turbulent diffusivity was considered to be constant at various depths and seasons, and the vertical component of current velocity in the ocean was not taken into account. ( Miyake and others-2 and Sereda-4).

Maturally only qualitative values were received as a result, but they showed much higher intensity of process of mixing in the ocean, than it followed the determination of age by radiocarbon.

The quantitative discription of process of spreading of radioactive contamination with stationary flow on the surface layer of the ocean( which shows contents of radioactive fall-out ) is given in this paper .For the analytical solution the authors had to schematize the problem showing ocean as a three-layered model with various laws of turbulent diffusivity distribution and vertical compoundants of current velocities in every layer .Till now quite sufficient scheme of spreading of radioactive contamination in the ocean was not received, that is why the following wor was performed by us:

1.During the r/v " Hikhail Lomonoecv " craises distribution of aplinter products, deposited on the ocean surface from the atmosphere has been studied.

2. The vertical turbulent diffusivity values and vertical components of the average current velocities were calculated by technique gives to the pener by 4.3. Folesnikov.

These calculations were based on measurement data of three fields—velocity of current, temperature and salinity, also obtained during expeditions aboard the r/v "X.Logonosov ".

3. The problem of finding arreading rate of radioactive contamination in the ocean can solved with the help of electronic computer . This solution was besed on the obtained turbulent diffusivity data and current velocity data not considering any simplifying suppositions.

Fresent paper is devoted to basic results of this work.

# 7 2.THE P/W " MINHAIL LONGHOSCY " OBJERVATIONAL DATA .

Curing recent years constant determinations of radionuclide contents in the ocean were performed about the research vessel of the Carine Hydrophysical Institute of the Academy of Ciences of the UKCO. Investigations were carried out both for surface watersand deep waters. The obtained samples were processed according to standard ther determination of radioactive or concentration by daughter ittrium-90. Carbonate precipitation of or-90 from 100-130 litre samples was made directly during expedition and received concentrated product was further prospected under stationary conditions.

Radio-chemical processing of concentrated material was in a large measure performed by scientific group under the leadership of professor V.P. Shvedov and candidate of Physical and Mathematical Science L. J. Godeonov whom authors are grateful . It was considered that tents of stable isotope of Sr with salimity 39.3% are 13 mg/ ag . It is natural that it warled depending on calinity and conditional density . As denoted contents of stable isotope are not surficient a portion of none radio active \$r- as salt SrCl2 - was added 0: the sample for the reliable precipitation with carrier. 750 g of warr ate of ammonium and 1200 g of anhydrous codium carbonate was added to the solution as a poiser in order to prevent magnesium precipitation. Sr output was controlled by flamephotometric method . Then Sr precipitated as strontium nitrate. Sediment of strontium nitrate was centrifuged, and radioactive Sr contents were determined by daughter Yttrium. The wast areas of the Atlantic Ocean ( 40-50 N to 10 S ) were studied . Considerable penetration of radioactive Sr can be traced at maximum depths , where determinations of this radioactive isotope were made . And it is characteristic of all measurements made .. Radio active Sr concentrations oscillated in different areas from 0.05 ( region of rise of deep water near Dakar ) to O. 13 pico-curie . That is in good agreement with data published previously. Lowering of activity to 0,02 o 0.05 pico-ourie can be noted in thermocline( See fag. 2). 1000 m lower to thermocline no abatement of specific actiwity is noticed -

§3.Calculation of penetration of radioactive contamination to depth.

The problem of vertical penetration of madicactive material

C concentration in the ocean can be given in the following equation:

$$\frac{\partial c}{\partial \tau} + W(z,\tau) \frac{\partial c}{\partial z} = \frac{\partial}{\partial z} \left( \partial(z,\tau) \frac{\partial c}{\partial z} \right) - \lambda c, /1/$$

with boundary conditions

$$C = C_0(\tau) \quad \text{with } z = 0 \qquad /2/2$$

$$\sum \frac{\partial C}{\partial z} = 0 \quad \text{with } z = 0 \qquad /3/2$$

where T is time, in vertical co-ordinate (axis = 0 = directed vertically down, it originates from the ocean surface . \* (z,T) - is vertical component of the average velocity of current .D (z,T) - is turbulent diffusivity. The is half-life of radiosctive material, H - ocean depth,

$$\lambda = \frac{0,693}{T_{1/2}}$$

Function Co (T), lineriting value of radioactive isotope concentration on the at lace was built using data obtained aboard the r/v "" all component " by measuring the weal contents of concentrations of radioactive material in the surface layer.

wand a values were computed using the actual data of measurements of our rent velocity, temperature and call - nity coursed out on board the r/v " h. homomosov " and also Fuglister 's data atlas ( 7 ) by technique given in the paper ( 6 ) The obtained " and D values were analysed and generalized by us and 'profiles of change of these values were drawn in accordance with depth and seasons ( for fig.1 ). Seasonal run of w and D variation with depth in the scheme accepted by us , corresponds to the following Triuss of thermooline depth according to months of the year;

XII. I. II / minter /-200 m.

III. IV. V / Opring / -60 m.

VI. VII. VIII / Summer /- 90 m.

XI. X. IX / Autumn / -150 m.

Paving determined  $C_0$ , E and  $\Psi$  aspect, the solution of problems /1/ — /3 / was made by electronic computer (by numerical method).

Computation involved period of time from 1954 to 1964. An example of such calculation of spreading of Sr-90 concentration \( \frac{7}{20} \) years / is given in fig.2; curve 1). This distribution of 3r-90 concentration with depth is referred to 4-th quarter of 1962. Data of 3r-90 concentration obtained during the r/v " Wikhail Lomonomov "expedition can be also referred to this period. They are given in fig.2. Theoretical and actual 6 values for 3r-90 are in good agreement to depth of 1000 m.

An assential effect of measonal variation of thermocline depth on ponetration of radioactive material from surface to deep water layers was revealed. It is known that minimam 0 values are found in lower border of thermocline . But directly under it ascending notions of water are tracad . Under such conditions blocking layer arises. Femateration of concentration lower to this layer is callicult of accessors to the accessor accessors and the accessors accessors conjugation of metorial in levering together with it, and as distributed nearly uniformly over it . After the rice es delma latrotes to antigration of material ( which was lowered together with it ) is found lower to it under conditions of marked values of turbulent diffucivity by the purtical line of Under these conditions the com sidered part of the material penetrates farther to deep waster layers without tariablons of thermeeline depth affect like a pluice, when consentration of radioastive material lo forle as four. Algeb of reyal scall and an effect. thomselian Clucications is of great importance in exchange of operation perspection balases surface inyer and deep wedday layur . To order to estimate the uffect of ecceimal washead on the same day the penetration of radio estive made this to propin an acculated it under the same conditions as for ourse to in fig. 2. But I end 9 ere independent of that, that is for the permanent depth of thermodilme .

The result of this calculation is shown as curve 2 in fig.2. As it is given in fig.2 specific activity values with variation of thermocline on the surface are twice less than at fixed profiles D and W.

## § 4. Conclusion.

Good agreement between actual and calculated concentrations shows the correctness of choice of initial suppositions. This circumstance allows us to detail a number of peculiarities of process of vertical epreading of radio active material in the ocean. In particulat, analysis of the given solution indicates that depth of penetration of radioactive isothess increases with the growth of half. life of radioactive isotope. Then we succeded in finding mechanism of spreading of isotope concentration in sur face layer caused by seasonal variations of thermocline depth, which creates the blocking layer. As it turned out the vertical transference of thermocline depth effects like a sluice letting radioactive material from surface layer in to deep-water layers. All this leads to much higher intensity of exchange than at permanent depth of thermocline. As a result the performed calculation fulfilled for ten year period indicates that during this period radioactive material from surface layer reached the bottom to depth of 5000 m, and consequently, it has much higher rates of spreading than according to "age".

DOCTOR OF PHYSICS AND MATHEMATICS

PROFESSOR

Skolejnikan/A.G. HOLESNIKOV/

CANDIDATE OF PHYSICS AND MATHEMATICS

Whelyaer / Porange/ 14. I. BELYAEY/

CANDIDATE OF PHYSICS AND MATHEMATICS

/B.A.NELEPO/

Pahelepo

### LITERATURE.

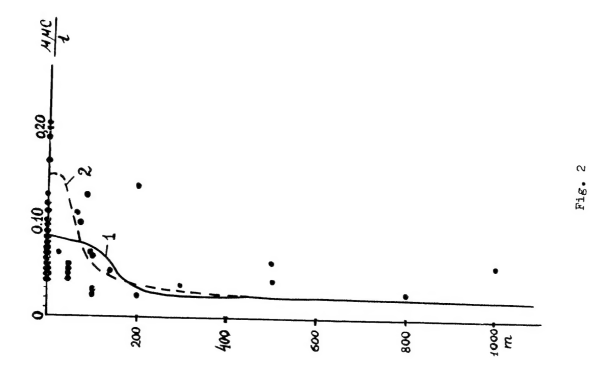
- 1.Bowen.V.T and Sugihara T.T.Larine geochemical studies with fall-out radioisotopes, Proc. Interm. Conf. Feaceful Uses At. Energy, 2nd , Geneva, 18, 434 , 1958 .
- 2.Miyake, Y. Saruhashi, Y. Katsuragi and T. Kanezawa. Penetration of Sr-90 and Cs-137 in deep layers of the Pacific and vertical diffusion rate of deep water .J. Radiation Res. 3.141-147.1962.
- 3.Rocco G.G., Proceder W.S. The vertical distribution of Cesium-137 indestrontium 90 in the oceans. J. of Geophysical Research.v.68. No.15.4501.
- 4. Sereda G.A. Contemination of seas and occars with artificial radioactive material.

  Problems of Nuclear Seteorology. Constantialat, 1962.
- 5.A.G.Koleanikov, A.A.Pivovarov and Z.G. Ivanova, On colculation of Rate of Radioactivity Spreading in Bepth of Gean. Abstracts of Symposium Papers, Y. Facific Science Congress, Honolulu, Hewaii, 346, 1961.
- 6.A.G. Kolesnikov. On the use of data of oxygen distribution for determination of vertical exchange in the ocean.

  Okeanology, v. 711. No 2,1965.
- 7. Euglister F.C.. Atlantic ocean atlas of temperature and salinity profiles and data from the International Geophysical Year of 1957-1958, USA. Boston, 1960.

# INCURIPTIONS TO FIGURES.

- Fig.1. Seasonal run of D and W variation with depth.
- Fig. 2. Distribution of Sr -90 concentration with depth at changing depth of thermosline (curve -1) and at permanent depth of thermosline (curve -2)



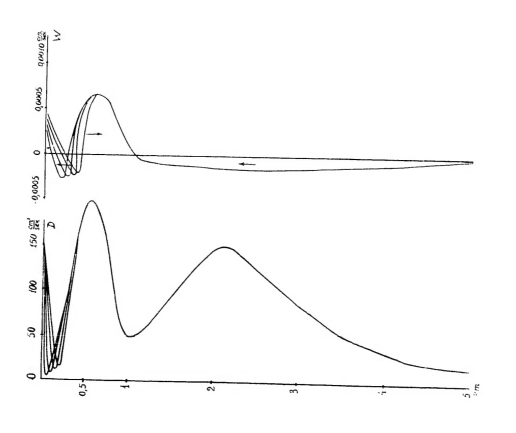


Fig. 1

766